PATENT SPECIFICATION

(11)1 409 009

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(21) Application No. 59232/73

(22) Filed 20 Dec. 1973 (31) Convention Application No. 127999/72

(32) Filed 20 D c. 1972 in

(33) Japan (JA)

(44) Complete Specification published 8 Oct. 1975

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G2C 212 242 26Y 304 30X 321 326 362 371 C19E2A C19HX

C2C 1452 20Y 213 247 250 252 25Y 292 29Y 30Y 366 367 37X 628 67Y 69Y 77Y 798 79Y CM ZG



(54) THERMALLY DEVELOPABLE LIGHT-SENSITIVE MATERIAL

(71) We, FUJI PHOTO FILM CO., LTD., a Japanese Company, of No. 210, Nakanuma, Minami/Ashigara-Shi, Kanagawa, Japan, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following

The present invention relates to a heat-developable light-sensitive material. Heretofore, photographic methods using silver halide have been widely practiced, since silver halides provide and the silver halides are silver halides sil

ERRATA

SPECIFICATION No. 1,409,009

Page 7, line 10, for monosulphate read monosulphonate

Page 12, line 23, after (C) insert was
Page 13, line 25, for 2-(p-dimethylaminostryryl) read 2-(p-dimethylaminostyryl)

THE PATENT OFFICE 1st December, 1975

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charge, or saccharm, or or benzotriazole, and a catalytic amount of a silver halide. The present invention relates to this recent proposal.

Heat-developable light-sensitive materials so far proposed are unfortunately insufficiently light-sensitive. It is however known that merocyanine dyes, the socalled colour sensitizers for silver halide emulsions, can also sensitize heatdevelopable light-sensitive materials.

Nonetheless, not all colour sensitizing agents effective for silver halide emulsions are necessarily effective for a heat-developable light-sensitive material.

We have discovered that a quinoline compound represented by the general formula (I) hereinafter described and certain derivatives thereof show good sensitizing effects for certain heat-developable light-sensitive materials.

Thus the present invention consists in a heat-developable light-sensitive material comprising a support bearing:

(a) an organic silver salt,

(b) a light-sensitive silver halide, or a compound capable of forming a light-

SEE ERRATA SLIP ATTACHED

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The present invention relates to a heat-developable light-sensitive material. Heretofore, photographic methods using silver halide have been widely practiced, since silver halides provide excellent sensitivity and gradation compared with electrophotographic methods or diazo-type photographic methods.

However, a silver halide light-sensitive material which is used in this method must, after imagewise exposure, be developed and then subjected to stopping, fixing, washing and stabilizing, in order to prevent the developed image from discolouring or fading and also to prevent the non-developed area (hereinafter referred to as "background") from blackening. Much time and labour are incurred and operators are exposed to danger from handling the necessary chemicals, and their hands and clothes may become stained.

Therefore, it would be desirable to process silver halide material in a dry state such that the processed images could be maintained stable.

Various proposals have been made to this end, for instance a combined developing and fixing method described in U.S. Patent 2,875,048, British Patent No. 954,453 and Federal German Patent No. 1,163,142, and a proposal to substitute dry processing operations for the wet processing operations in the conventional silver halide photographic method, as described in Federal German Patent No. 1,174,159 and British Patent Nos. 943,476 and 951,644. A recent proposal, as described in Japanese Patent Publication No. 22185/70, U.S. Patents 3,152,904, 3,457,075, 3,635,719 and 3,645,739 and British Patent No. 1,205,000, is to use a heat-developable light-sensitive material containing a light-sensitive element comprising mainly a silver salt such as a long-chain carboxylic acid (e.g. silver behenate), of saccharin, or of benzotriazole, and a catalytic amount of a silver

halide. The present invention relates to this recent proposal. Heat-developable light-sensitive materials so far proposed are unfortunately insufficiently light-sensitive. It is however known that merocyanine dyes, the socalled colour sensitizers for silver halide emulsions, can also sensitize heatdevelopable light-sensitive materials.

Nonetheless, not all colour sensitizing agents effective for silver halide emulsions are necessarily effective for a heat-developable light-sensitive material.

We have discovered that a quinoline compound represented by the general formula (I) hereinafter described and certain derivatives thereof show good sensitizing effects for certain heat-developable light-sensitive materials.

Thus the present invention consists in a heat-developable light-sensitive material comprising a support bearing:
(a) an organic silver salt,

(b) a light-sensitive silver halide, or a compound capable of forming a light-

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sensitive silver halide by reaction with said organic silver salt (a), in an amount effective to catalyze the reaction of compounds (a) and (c),

(c) a reducing agent capable of reducing to elemental silver the organic silver salt (a) when the material is being heated in the presence of the component (b) which has been exposed to light,

(d) a bind r, which may be omitted if the organic silver salt (a) and/or the reducing agent (c) act as a binder,

(e) a quinoline compound represented by the following general formula (I):

$$N = \frac{CH - CH}{\underline{a}} C - \frac{(X - Y)}{\underline{n}} D \quad (I)$$

10 R1 and R2 being each independently an alkyl group having from 1 to 4 carbon atoms or a benzyl group; X and Y are each independently =C—or =N—; n is 1 or 2; a is 0 or 1; and Q represents the atoms necessary to complete a quinoline nucleus which may be further substituted; or

(e) may be a quinoline ring N-oxide derived from said quinoline compound or a

quinolinium salt derived from said quinoline compound R^1 and R^2 are preferably methyl, ethyl or propyl such as n-propyl. R^1 and R^2 can be the same or different. Also, the quinoline nucleus in the general formula can be substituted, such as by an alkyl group having 1 to 4 carbon atoms, such as methyl, ethyl, isopropyl or isobutyl (any of which may be substituted with a halogen atom or a hydroxyl group, thus being e.g. hydroxyethyl or β -chloroethyl), a phenyl group, an alkoxy group having 1 to 4 carbon atoms such as methoxy, ethoxy, propoxy or butoxy, a halogen such as a fluorine, chlorine, bromine or iodine, a dialkylamino group in which the alkyl moiety may have I to 4 carbon atoms (e.g. methyl, ethyl, propyl, isopropyl, butyl or t-butyl), an alkoxycarbonyl group having I to 4 carbon in the alkoxy moiety (such as a methoxycarbonyl or ethoxycarbonyl group) and $-(X=Y)_aD$, X, Y, n and D being as hereinbefore defined in connection with the general formula (I).

Now, specific examples of the compounds represented by the general formula (I), quinoline ring N-oxides thereof and quinolinium salts thereof, useful as the sensitizing agent in the present invention, are given below:

(1) 4-(p-dimethylaminostyryl)quinoline

(2) 4-(4-p-dimethylaminophenyl-1,3-butadienyl)quinoline

(3) 2-(4-dimethylaminostyryl)quinoline

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(4) 2-(p-dimethylaminobenzylidene)aminoquinoline

(5) 4-(p-dimethylaminophenylimino)methylquinoline

(6) 4-(p-dimethylaminophenylazo)quinoline

(7) 4-I(3-p-dimethylaminophenyl)-2-propenylidenel-aminoquinoline

(8) 6-chloro-2-(p-dimethylaminostyryl)quinoline

(9) 6-methoxy-2-(p-dimethylaminostyryl)quinoline

(10) 8-fluoro-4-(p-dimethylaminostyryl)quinoline

15 (11) 4-(p-dipropylaminostyryl)quinoline

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(12) 4-(p-methylbenzylaminostyryl)quinoline

(13) 4-(p-dimethylaminostyryl)quinoline-1-oxide

5 (14) 4-(p-diethylaminophenylimino)methylquinoline-1-oxide

C2H5

(15) 2-[4-(p-dimethylaminophenyl)-1,3-butadienyl]quinoline-1-oxide

(16) 4-(p-dimethylaminostyrylquinolinium) ethyl iodide

(17) 4-(3,4-dioxymethylenephenylethylenyl)quinolinium ethyl iodide

(18) 2,4-bis(p-dimethylaminostyryl)quinolinium ethyl iodide

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(19) 4-(p-dimethylaminostyryl)quinolinium 1,2-dihydroxypropyl tosylate

(20) 4-(p-dimethylaminostyryl)quinolinium n-heptyl iodide

(21) 2-(p-dimethylaminostyryl)-6-dimethylaminoquinolinium ethyl iodide

CH₃

$$N$$
 $CH = CH$
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3

(22) 4-[4-(p-dimethylaminophenyl)-1,3-butadienyl]quinolinium n-heptyl iodide

$$n-C7H_{15}$$
 N
 $CH=CH$
 $CH=CH$
 $CH=CH$

(23) 2-[4-(p-dimethylaminophenyl)-1,3-butadienyl]quinolinium ethyl bromide

These compounds can be used alone or in combinations of two or more.

Compounds (1), (17), (20), (21), and (22) are particularly preferred.

The compound effective as the sensitizing agent in material according to the present invention (component (e)) is suitably added in an amount of from 10⁻⁶ mol to 10⁻² mol, preferably 10⁻³ mol to 10⁻³ mol, per 1 mol of the organic silver salt (a), the amount chosen depending for example upon the kind of compound used, the kind of organic silver salt, the kind of silver halide, the kind of reducing agent and the processing temperature. If the amount of the component (e) exceeds 10-2 mol per mol of organic silver salt, and the component (e) is one giving rise to considerable density of colour, the degree of colouration of the light-sensitive layer can become too great, which is not desirable for some uses.

Suitable examples of organic silver salts which can be used as component (a) in the present invention are the silver salts of organic compounds having an imino group or a mercapto group, and the silver salts of organic (preferably carboxylic) acids, as disclosed in U.S. Patent 3,457,075. Silver salts of organic acids having 10

or more carbon atoms are preferable.

	1,405,005	6
	Specific examples of these silver salts include the silver salt of benzotriazole, of saccharin, of phthalazinone, of 3-mercapto-4-phenyl-1,2,4-triazole, of 4-hydroxy-6-methyl-1,3,3a,7-tetrazaindene, of 2-(S-ethylthioglycolamido)-benzothiazole, silver caprate, silver laurate, silver myristate, silver palmitate, silver stearate, silver behanne, silver adjects of the silver palmitate, silver sil	
5	Examples of component (b) in the present invention, capable of forming a light-sensitive silver halide by the reaction thereof with the organic silver salt (a), include hydrogen halide, ammonium halide or a motal belief	.5
10	being strontium, cadmium, zinc, tin, chromium, sodium, barium, iron, caesium, lanthanum, copper, calcium, nickel, magnesium, potassium, aluminium, antimony, gold, cobalt, mercury, lead, beryllium, lithium, manganese, gallium, indium, platinum, thallium or bismuth, and the halide being the chloride, bromide or lodide.	10
15	Other examples of component (b) are organic halogen compounds such as monochlorotriphenylmethane, monobromotriphenylmethane, 2-bromo-2-methyl-propane, 2-bromobutyric acid, 2-bromoethanol, dichlorobenzophenone, triiodomethane, tribromomethane and tetrabromomethane. Suitable silver halides for use as component (b) are silver chloride, silver bromoided silver halides for use as component (b) are silver chloride, silver bromoided.	15
20	bromoiodide, silver chlorobromide and silver iodide. These light-sensitive silver halides can be of coarse grains or of fine grains. Emulsions of extremely fine grains are particularly useful. Emulsions containing a light-sensitive silver halide (b) can be respected.	20
25	method, the double jet method (e.g. a Lippmann's emulsion), or are an ammoniacal emulsion, a thiocyanate- or thioether-ripened emulsion, or are as described in, e.g. U.S. Patents 2,222,264, 3,320,069 and 3,271,157. Silver halides (b) used in material according to the recent investigation.	25
30	selenium compounds, gold compounds, platinum compounds, palladium compounds, or combinations thereof. Suitable procedures for chemical sensitization are described in, e.g. U.S. Patent 2,623,499, 2,399,083, 3,297,447 and 3,297,446.	30
35	The compounds which can be employed as component (b) may be used independently or as a combination of two or more. This component (b) is suitably added in an amount of from 0.001 mol to 0.5 mol per 1 mol of the organic silver salt, component (a). If the amount of the component (b) is less than 0.001 mol per mol of (a), the sensitivity is reduced, while if it exceeds 0.5 mol per mol of (a), the	35
40	leaving the material open to light, diminishing the contrast between the image areas and non-image areas. The reducing agent which is used as component (c) in the present invention must be capable of reducing the organic silver salt (a) to form elemental silver.	40
45	component (b), which is a silver halide catalyst or can form such. The best reducing agent for a particular material depends upon the organic silver salt used. For example, when using a silver salt of a higher fatty acid such as silver behenate, or using the silver salt of benzatriazele, which are	45
50	difficult to reduce, comparatively strong reducing agents such as a bis-phenol (e.g. 4,4'-methylenebis-(3-methyl-5-t-butylphenol) or an ascorbic acid respectively are suitable. On the other hand, comparatively weak reducing agents such as a substituted phenol (e.g., p-phenylphenol) are suitable for silver salts which are comparatively easy to reduce, such as silver laurate. Examples of frequently useful	50
55	reducing agents are substituted phenols, substituted or unsubstituted bis-phenols, substituted or unsubstituted bis-phenols, substituted or unsubstituted bis-naphthols, di- or poly-hydroxynaphthalenes, di- or higher-polyhydroxynaphthalenes, di- or higher-polyhydroxynaphthalenes, hydroquinone monoethers, di-ethers, and tetra-ethers, ascorbic acid or derivatives thereof, 3-pyrazolidone, pyrazolin-5-ones, reducing saccharides (sugars), kojic acid and hinokitiol.	55
60	those where the substituted phenois, bisphenois, bisnaphthols and naphthols are those where the substituent groups are alkyl (C_1-C_8) , alkoxy (C_1-C_8) , phenyl, halogen, amino, alkyl substituted amino (C_1-C_8) , benzyl, hydroxyalkyl (C_1-C_8) , acetyl or nitro.	60
65	Most preferred of the hydroquinone ethers are the monoalkyl (C_1-C_8) ethers, the monoaralkyl ethers and the monoaryl ethers.	65

<u> </u>	1,409,009	8
	from 0.2 to 3 g of silver per square metre of the support, preferably 0.4 to 2 g of silver per square metre.	
5	An antistatic layer or an electroconductive layer can be provided on the heat- developable light-sensitive material of the present invention if desired. Also, an antihalation substance or an antihalati n dye can be incorporated therein, as	5
	If desired, a matting agent such as starch, titanium dioxide, zinc oxide or silica, as well as a fluorescent brightening agent such as a stilbene, triggine, oxygole	J
10	or coumarin can be additionally incorporated in the material. The components (a) to (e) and any additives can be applied on a support, to	10
10	various coating methods including immersing methods, air-knife coating methods, curtain-coating methods and extrusion coating methods using a hopper as	10
15	described in U.S. Patent 2,681,294. If desired, two or more layers can be applied at the same time. The components (a) to (e) and any additives may be present in a	15
13	single layer or some may be applied as separate layers. Some optically sensitizing dyes used for silver halide emulsions can increase	15
	the light sensitivity of the heat-developable light-sensitive material of the present invention. Useful optical sensitizers include the cyanine dyes and the merocyanine	
20	dyes, as disclosed in U.S. Patents 3,457,075 and 3,761,279. The amount used of the	20
	cyanine or merocyanine dyes is preferably from 10^{-6} mol to 10^{-2} mol per 1 mol of the organic silver salt (a). In order to enhance the transparency of the thermally developable light-	
25	sensitive layer, to increase the image density and to improve the shelf-life of the	
25	material, a top-coating polymer layer, preferably 1 to 20 microns thick, can be	25
	provided on the light-sensitive layer. Suitable top-coating polymers include polyvinyl chloride, polyvinyl acetate, vinyl chloridevinyl acetate copolymers,	
	polyvinyi butyral, polysturene, polymethyl methacrylate, polyurethane rubbers	
30	xylene resins, benzyl cellulose, ethyl cellulose, cellulose acetate butyrate, cellulose acetate, polyvinylidene chloride, chlorinated polypropylene, polyvinyl	30
	pyrrolidone, cellulose propionate, polyvinyl formal, cellulose acetate phthalate	30
	polycarbonates and cellulose acetate propionate.	
	Heat-developable light-sensitive material according to the invention can be developed, after being exposed to irradiation in the ultraviolet and/or visible range	
35	from a xenon lamp, tungsten lamp, mercury lamp, or other light source, by merely	35
	heating the light-sensitive material. Heating temperatures of 100 to 160°C, more preferably 110 to 140°C, are suitable for such development. Higher or lower	
	temperatures can be selected within the above-described range, shortening or	
40	prolonging the developing time respectively. The developing time is usually from 1	40
40	second to 60 seconds. The material can be heated, for example, by being brought into contact with a heating plate or the like or with a heated drum, or, in some	40
	cases, by being passed through heated space, or by using high-frequency induction	
	heating or a laser beam. The present invention will now be illustrated in greater detail by reference to	
45	the following examples of preferred embodiments. Unless otherwise indicated, all	45
	parts, percentages and ratios are by weight.	
	EXAMPLE 1.	
	3.4 Grams of behenic acid was dissolved in 100 ml of toluene at 60°C, and the solution temperature was held to 60°C. Then, 100 ml of a dilute nitric acid aqueous	
50	solution of a pH of 2.0 (at 25°C) was mixed thereinto with stirring. This solution	50
	mixture was maintained at 60°C, and to this was added, while continuing the	
	stirring, an aqueous solution prepared by adding aqueous ammonia to about 80 ml of an aqueous solution containing 1.7 g of silver nitrate to form a silver ammonium	
	complex salt and making the total amount 100 ml with water. Thus, a dispersion	
55	containing fine crystals of silver behenate was obtained. Upon leaving this dispersion for 20 minutes at room temperature (about 20—30°C), it separated into	55
	an aqueous phase and a toluene phase.	
•	First, the aqueous phase was removed. Then, 400 ml of fresh water was added	
60	thereto for washing followed by decanting. This procedure was repeated 3 times. Then, 400 ml of water was added thereto and silver behenate was collected by	60
	centrifuging. Thus, 4 g of rod-shaped silver behenate crystals of about 1 micron in	•
	length and about 0.05 micron in width were obtained, 2.5 Grams of the thus	
	obtained silver behenate was added to 20 ml of an isopropyl alcohol solution containing 2 g of polyvinyl butyral and was subjected to ball milling for 1 hour to	
	O O 1 - 7 - 7 - 2-7	

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5	prepare a polymer dispersion. To 20 ml of this polymer dispersion of the silver salt were added the following components to prepare a thermally developable light-sensitive composition. This composition was then applied to a polyethylene terephthalate film support in a silver amount of 1.5 g per 1m ² of the support to prepare a thermally developable Light-sensitive Material (A).		
	Ammonium bromide (2.5% by weight methanol solution) 1 ml		
	Compound (1) (0.025°; by weight 2-methoxy-ethanol solution)		
10	2,2'-Methylene <i>bis</i> (6- <i>t</i> -butyl-4-methyl-phenol) (25°, by weight solution in 2-methoxy-ethanol) 3 ml	10	
	Phthalazinone (2.5% by weight solution in 2-methoxy-ethanol) 1 ml		
15	Tetrachlorophthalic anhydride (0.6% by weight methanol solution)	15	
20	Separately, for the purpose of comparison, there were prepared in a similar manner a thermally developable Light-sensitive Material (B) not containing developable Light-sensitive material (C) containing as a sensitizing dye a in the same amount. Merocyanine Dye:		
	- 7		

On each of these Light-sensitive Materials (A) to (C) was top-coated a 15% by weight tetrahydrofuran solution of vinyl chloride-vinyl acetate copolymer (comprising 95% by weight of vinyl chloride and 5% by weight of vinyl acetate) in a Each of the 1 light const.

Each of these Light-sensitive Materials (A) to (C) was exposed to 240,000 lux-sec from a tungsten light source through an optical wedge. Thereafter, the materials were heated at 120°C for 10 seconds to develop, then the resulting blackened density was measured.

The reciprocal number of exposure amount necessary to provide a density higher than fog (transmittance at the unexposed area upon heating) by 0.1 was selected to express the sensitivity. The results obtained are shown below wherein the relative sensitivity of the Light-sensitive Material (B) was taken as 100.

		- (-) " and tartest the 100.		
	Light-sensitive Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)	
Relative Sensitivity	20,000	100	500	

From the above-described results, it can be seen that the sensitizing agent of the present invention exhibits an outstanding sensitizing effect.

EXAMPLE 2. The same procedures as described in Example 1 were conducted except for using both Compound (1) and the merocyanine dye used in Example 1 in the same amounts in place of Compound (1). The results obtained are shown 40

	Light-sensitive Material (A) (containing compound (1) and the merocyanine dye	Light-sensitive Material (B) (containing neither Compound (I) nor merocyanine dye)	Light-sensitive Material (C) (containing the merocyanine dye)
Relative Sensitivity	25,000	100	500
It can be seen fr and the merocyanine	rom the results, that dye served to obtain EXAMPI	an even more exc	e of Compound (1) cellent effect.
The same proced for using Compound (dures as described in (21) in place of Com	n Example I were	conducted except ame amount. Thus,
	Light-sensitive Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)
Relative Sensitivity	5,200	100	500
The same effect	as in Example 1 was	observed.	-
The same proceed for using Compound of follows.	EXAMPI dures as described i (7) in place of Comp	n Example i were	conducted except ults obtained are as
	Light-sensitive	li-baisi	I tale a tet
	Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)
Relative Sensitivity		Material (B)	
maintaining the solution was added thereto wit aqueous solution (concontaining 8.5 g of silve the lauric acid with the lauric no length and silver salt was washed polyvinyl butyral and laurate and dispersed usalt.	Material (A) 3,800 EXAMPI ic acid was dissolved on at 10°C, 100 ml o th stirring. Then, whi oled to 0°C) of a s er nitrate was added e silver ion. Thus, roc d about 0.05 micron successively, with wa 20 ml of isopropyl using a ball mill to pre olymer dispersion of e a thermally develow was applied to a polyo per 1 m² of the supp	Material (B) 100 LE 5. In 100 ml of butyle f aqueous nitric actile continuing the silver nitrate ammon thereto over a 1 mill-shaped silver laurain width were obtained alcohol were added the silver salt were pable light-sensitive thylene terephthal	Material (C) 500 I acetate and, while id (pH at 25°C: 2.0) stirring, 50 ml of an onium complex salt nute period to react ate crystals of about ained. The resulting Thereafter, 3.0 g of id to 2.7 g of silver persion of the silver added the following e composition. The ate film support in a
11 Grams of laur maintaining the solution was added thereto with aqueous solution (containing 8.5 g of silvethe lauric acid with the lauric acid with the lauric acid with the lauric acid with the lauric and laurate and dispersed usalt. To 20 ml of this percomponents to prepar resulting composition silver amount of 1.7 g light-sensitive materia	EXAMPI ic acid was dissolved on at 10°C, 100 ml o th stirring. Then, whi oled to 0°C) of a s er nitrate was added e silver ion. Thus, roc d about 0.05 micron successively, with wa 20 ml of isopropyl using a ball mill to pre olymer dispersion of e a thermally developer was applied to a polyc per 1 m² of the supp I (A).	Material (B) 100 LE 5. I in 100 ml of buty f aqueous nitric acile continuing the silver nitrate ammon thereto over a 1 mill-shaped silver laurin width were obtained at er and methanol, alcohol were addeepare a polymer distinct the silver salt were expable light-sensitive thylene terephthal ort to prepare a the	Material (C) 500 I acetate and, while id (pH at 25°C: 2.0) stirring, 50 ml of an onium complex salt nute period to react ate crystals of about ained. The resulting Thereafter, 3.0 g of id to 2.7 g of silver persion of the silver added the following e composition. The ate film support in a
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11 Grams of laur maintaining the solution was added thereto with aqueous solution (concontaining 8.5 g of silve the lauric acid with lauric acid was washed polyvinyl butyral and laurate and dispersed to salt. To 20 ml of this promponents to prepare resulting composition silver amount of 1.7 g light-sensitive materia Ammonium bro (2.5°, by weigh Compound (20) (0.025°, by weigh p-Phenylphenol	EXAMPI ic acid was dissolved on at 10°C, 100 ml o th stirring. Then, who led to 0°C) of a s er nitrate was added e silver ion. Thus, roo d about 0.05 micron successively, with wa 20 ml of isopropyl using a ball mill to pro olymer dispersion of e a thermally develo was applied to a polyo per 1 m² of the supp I (A). omide t methanol solution) ght 2-methoxy-ethan	Material (B) 100 LE 5. In 100 ml of buty f aqueous nitric acille continuing the silver nitrate ammon thereto over a l mileshaped silver laurain width were obtained alcohol were addespare a polymer distinct the silver salt were pable light-sensitive thylene terephthal ort to prepare a the solution)	Material (C) 500 l acetate and, while id (pH at 25°C: 2.0) stirring, 50 ml of an onium complex salt nute period to react at a crystals of about ained. The resulting Thereafter, 3.0 g of id to 2.7 g of silver persion of the silver added the following e composition. The ate film support in a crmally developable

 	1,409,0	09		11
Tetrachlorophtha (0.6% by weight r	lic anhydride nethanol solution)		1 ml	
Separately, for the purposes of comparison, there were prepared in a similar manner a thermally developable Light-sensitive Material (B) not containing Compound (20) and a thermally developable Light-sensitive Material (C) containing as a sensitizing dye a merocyanine dye (the same dye as used in Example 1) in place of Compound (20) in the same amount. One each of these Light-sensitive Materials (A) to (C) was top-coated a 15% by weight tetrahydrofuran solution of vinylarchloride-vinyl acetate copolymer (comprising 95% by weight of vinyl chloride and 5% by weight of vinyl acetate) in a				
Each of these three tungsten light source; thr 240,000 lux sec. Thereaft develop, and then the re The relative sensitiv The results thus obtaine	er, the materials vesulting blackened	vere heated at 120° density was measu	C for 10 seconds to	.1
	Light-sensitive Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)	
Relative Sensitivity	4,700	100	500	
From the above-des according to the present	cribed results, it invention shows	can be seen that than outstanding effe	ne sensitizing agent	:
The same procedure using Compound (18) in results thus obtained are	given in the follo	Example 5 were co ound (20) in the s wing table. Light-sensitive	Light-sensitive	
Relative Sensitivity	Material (A)	Material (B)	Material (C)	
The same effect as	1,200	100	500	
The same procedure using Compound (4) in p thus obtained are given	EXAMPL es as described in lace of Compound in the following ta	Example 5 were co	nducted except for mount. The results	
	Light-sensitive Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)	
Relative Sensitivity	1,400	100	500	
The same effect as i	n Example 5 was	observed.		
6 Grams of benzotri:	EXAMPL	E 8.		
and the solution was cool an aqueous solution preparation (pH temperature of 3°C. Thus benzotriazole was obtain room temperature, it sephase. First, the aqueous ph thereto for washing follow.	led to -15°C. To the total series of the total	his solution was add 8.5 g of silver nitrate and the mixture value of the crystals this dispersion for a company the compan	e in 100 ml of dilute was adjusted to a of the silver salt of 20 minutes at an an isoamyl acetate is water was added	

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obtained silver salt of benzotriazole was added to 40 ml of an isopropyl alcohol solution containing 4 g of polyvinyl butyral and was subjected to ball milling for 4 hours to prepare a polymer dispersion of the silver salt. To 40 ml of this polymer dispersion of the silver salt were added the following components to prepare a thermally developable light-sensitive composition. This composition was then applied to a polyethylene terephthalate film support in a silver amount of 1.2 g per 1 m ² of the support to prepare a thermally developable Light-sensitive Material (A).					
	Ammonium Iodi (8.5° by weight	de methanol solution)	1 ml	
	Solution contain Monopalmitate a in 10 ml of 2-me	ing 2 g of Ascorbi and 2 g of Ascorb thoxy-ethanol	c Acid ic Acid Dipalmitate	10 ml e	
	Compound (8) (0.2°, by weight	2-methoxy-ethano	l solution)	l ml	
	N-Ethyl-N'-dode (2.5°, by weight	cylurea 2-methoxy-ethano	solution)	2 ml	
	Compound (8) and a containing as a sensitizing Example 1) in place of these I weight tetrahydrofuran (comprising 95% by weight thickness of 8 micro	thermally development of the thermally development (8) in clight-sensitive Material solution of a vight of vinyl chloride ons	lopable Light-sens nine dye (the same the same amount, terials (A) to (C) to and 5% by weight (S. (A) to (C) was and (C) to the same the same amount.	dye as described in op-coated a 15% by acetate copolymer of vinyl acetate) in a	
	light source through an lx sec. Thereafter, the mThen, the resulting black determined in the same obtained are tabulated to	naterials were heate kened density was manner as descri	inpart thereto an exect at 130°C for 30;	posure of 1,200,000 seconds to develop.	
	lx-sec. Thereafter, the m Then, the resulting black determined in the same obtained are tabulated b	naterials were heate kened density was manner as descri	inpart thereto an exect at 130°C for 30;	posure of 1,200,000 seconds to develop.	
	lx-sec. Thereafter, the m Then, the resulting black determined in the same	kened density was manner as describelow. Light-sensitive	ed at 130°C for 30 s measured. The relatibed in Example Light-sensitive	posure of 1,200,000 seconds to develop. ative sensitivity was 1. The results thus Light-sensitive	•
	lx-sec. Thereafter, the m Then, the resulting black determined in the same obtained are tabulated b	Light-sensitive Material (A) 450 escribed results	ed at 130°C for 30 s measured. The rela ribed in Example Light-sensitive Material (B) 100	posure of 1,200,000 seconds to develop. ative sensitivity was 1. The results thus Light-sensitive Material (C)	·
	lx-sec. Thereafter, the m Then, the resulting black determined in the same obtained are tabulated by Relative Sensitivity From the above-d	Light-sensitive Material (A) 450 escribed results, invention shows EXAMPI ures as described pound (3) in pla	Light-sensitive Material (B) 100 it can be seen of an outstanding effect. E 9. d in Example 8	Light-sensitive Material (C) 150 that the sensitizer ect.	•
	Ix-sec. Thereafter, the m Then, the resulting black determined in the same obtained are tabulated to the same obtained are tabulated to the same proced according to the present of the same proced except for using Committee to the same proced except for using Committee to the same procede to the same procede except for using Committee to the same obtained are tabulated to the same obtained to the same obtained are tabulated to the same obtained to the same obtain	Light-sensitive Material (A) 450 escribed results, invention shows EXAMPI ures as described pound (3) in pla	Light-sensitive Material (B) 100 it can be seen of an outstanding effect. E 9. d in Example 8	Light-sensitive Material (C) 150 that the sensitizer ect.	-

EXAMPLE 10.

The same procedures as described in Example 8 were conducted except for using Compound (13) in place of Compound (8) in the same amount. Thus, the results as given in the following table were obtained.

	Light-sensitive Material (A)	Light-sensitive Material (B)	Light-sensitive Material (C)
Relative Sensitivity	390	100	150
The same effect as	in Example 8 was	observed.	· · · · · · · · · · · · · · · · · · ·
WHAT WE CLAIM 1. A heat-developab (a) an organic silver (b) a light-sensitive s sensitive silver halide by effective to catalyze the (c) a reducing agent salt (a) when the materia which has been exposed (d) a binder, which reducing agent (c) act as (e) a quinoline comp	le light-sensitive me salt, silver halide, or a concept reaction with said reaction of compostable of reducing all is being heated it to light, may be omitted if a binder.	compound capable organic silver salt ounds (a) and (c), g to elemental silve in the presence of the organic silver	of forming a light- t (a), in an amount or the organic silver the component (b) salt (a) and/or the
	<u></u>	Ţ	
	і N == (сн− сн -}= -	C (x x y) D	(1)
	` ' <u>a</u>	'n	•
	R1		CH ₂
wherein-D 15-	-	or —	
	R ²		
R ¹ and R ² being each			u
carbon atoms or a ben or =N; n is 1 or 2; a is 0 o quinoline nucleus which n N-oxide derived from said said quinoline compound. 2. A light-sensitive m methyl, ethyl or n-propyl.	or 1; and Q represe nay be further subs quinoline compou	nts the atoms neces tituted; or (e) may l nd or a quinoliniun	lependently =C— sary to complete a be a quinoline ring a salt dervied from
3. A light-sensitive mais 4-(p-dimethylaminosty aminoquinolinium ethyliodide, 4-(3,4-dioxymethyldimethylaminophenyl)-1 3	aterial as claimed in yryl)quinoline, 2- iodide, 4-(p-dimet enephenylethyleny	hylaminostyryl)qui hylaminostyryl)qui l)quinolinium ethyl	iryryl)-6-dimethyl- nolinium n-heptyl liodide, or 4-[4-(p-
component (e) is present i silver salt (a).	material as claime in an amount of 10	ed in any precedin -6 to 10 ⁻² mol per n	ig claim, wherein nol of said organic
5. A light-sensitive me support bears a cyanine of 6. A light-sensitive ma said merocyanine dye is porganic silver salt (a).	iye or a merocyan iterial as claimed in	ine dye. Claim 6 wherein c	aid avanina dva a-
7. A light-sensitive ma organic silver salt (a) is a s compound containing an	inino group or a	anic carboxylic aci mercanto group	d or of an organic
o. A light-sensitive ma compound capable of form organic silver salt (a) is a h	aterial as claimed i ning a light-sensiti nydrogen halide, an bromide or jodide	in any preceding cl ve silver halide by ammonium halide	reaction with said or a metal halide,
9. A light-sensitive ma			

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5	potassium, aluminium, antimony, gold, cobalt, mercury, lead, beryllium, lithium, manganese, gallium, indium, platinum, thallium or bismuth. 10. A light-sensitive material as claimed in any of Claims 1 to 7, wherein said compound capable of forming a light-sensitive silver halide by reaction with said organic silver salt (a) is monochlorotriphenylmethane, mon bromotriphenylmethane, 2-bromo-2-methylpr pane, 2-bromo-butyric acid, 2-bromoethanol, dichlorobenzoph none, triiodomethane, tribromomethane or tetrabromomethane.	5
10	11. A light-sensitive material as claimed in any preceding claim, wherein said light-sensitive silver halide is silver chloride, silver bromide, silver bromoiodide, silver chlorobromoiodide, silver chlorobromide or silver iodide. 12. A light-sensitive material as claimed in any preceding claim, wherein said reducing agent (a) is a substituted as claimed in any preceding claim, wherein said	10
15	reducing agent (c) is a substituted phenol, a substituted or unsubstituted bisphenol, a substituted or unsubstituted in aphthol or bis-naphthol, a di- or poly-hydroxy-naphthalene, di- or higher- polyhydroxybenzene, di- or higher- polyhydroxy-naphthalene, a hydroquinone mono-, di-, tri- or tetra-ether, ascorbic acid or a derivative thereof, a 3-pyrazolidone, a pyrazolin-5-one, a reducing sugar, kojic acid or hinokitiol.	15
20	13. A heat-developable light-sensitive material as claimed in Claim 1 and substantially as hereinbefore described with reference to and as shown in any one of Examples 1 to 10. 14. A method of developing a light-sensitive material as claimed in claimed in the sensitive material as claimed in Claim 1 and substantially as sensitive material as claimed in Claim 1 and substantially as sensitive material as claimed in Claim 1 and substantially as hereinbefore described with reference to and as shown in any one of Examples 1 to 10.	20
25	the material to a temperature for a duration until a black image is obtained. 15. A method as claimed in Claim 14, wherein the temperature is from 100° to 16. A method as claimed in Claim 15, wherein the temperature is from 100° to	25
30	140°C. 17. A method as claimed in any of Claims 14 to 16, wherein the duration is from 1 to 60 seconds. 18. A material bearing a visible image developed by a process as claimed in any of Claims 14 to 17.	30

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